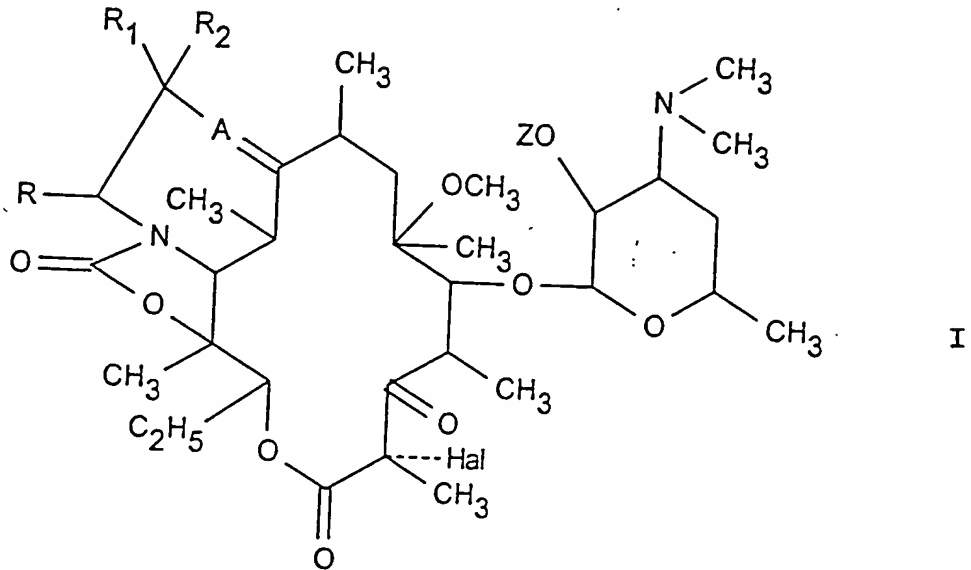


WHAT WE CLAIM IS:

1. A compound selected from the group consisting of a compound of
5 the formula



10
15
wherein A is nitrogen or N>O, R₁ and R₂ are individually selected from the group consisting of hydrogen and alkyl of 1 to 18 carbon atoms, R is selected from the group consisting of hydrogen and -(CH₂)_mOB, Hal is halogen, m and n are individually an integer
20
from 1 to 8, B is hydrogen or -C=Ar₂OR-(CH₂)_n-Ar, Ar is a mono- or polycyclic aryl or heteroaryl, Z is hydrogen or acyl of an organic carboxylic acid of up to 18 carbon atoms and its non-toxic, pharmaceutically acceptable acid addition salts.

25

2. A compound of claim 1 wherein R₁ and R₂ are hydrogen.

3. A compound of claim 1 wherein A is nitrogen.

4. A compound of claim 1 wherein Hal is fluorine.

5 5. A compound of claim 1 wherein R is hydrogen.

6. A compound of claim 1 wherein R is -CH₂OH.

7. A compound of claim 1 selected from the group consisting of
10 [3aS-(3aR*,4S*,7R*,9S*,10S*,11S*,13S*,15S*,15aS*)]-4-ethyl-7-
fluoro-3a,4,10,11,12,13,15,15a-octahydro-11-methoxy-
3a,7,9,11,13,15-hexamethyl-10-[(3,4,6-trideoxy-3-(dimethyl-amino)-
.beta.-D-xylo-hexopyranosyl)oxy]-14,1-(nitriloethano)-2H-
oxacyclotetradecino[4,3-d]oxazole-2,6,8(9H)-trione and
15 [3aS-(3aR*,4S*,7R*,9S*,10S*,11S*,13S*,15S*,15aS*,17R*)]-4-
ethyl-7-fluoro-3a,4,10,11,12,13,15,15a-octahydro-17-hydroxymethyl)-
11-methoxy-3a,7,9,11,13,15-hexamethyl-10-[(3-4,6-trideoxy-3-
(dimethylamino)-.beta.-D-xylohexopyranosyl)oxy]-14,1-
.✓ (nitriloethano)-2H-oxacyclotetradecino[4,3-d]oxazole-2,6,8(9H)-
20 trione.

8. An antibiotic composition comprising an antibiotically effective amount of a compound of claim 1 and an inert pharmaceutical carrier.

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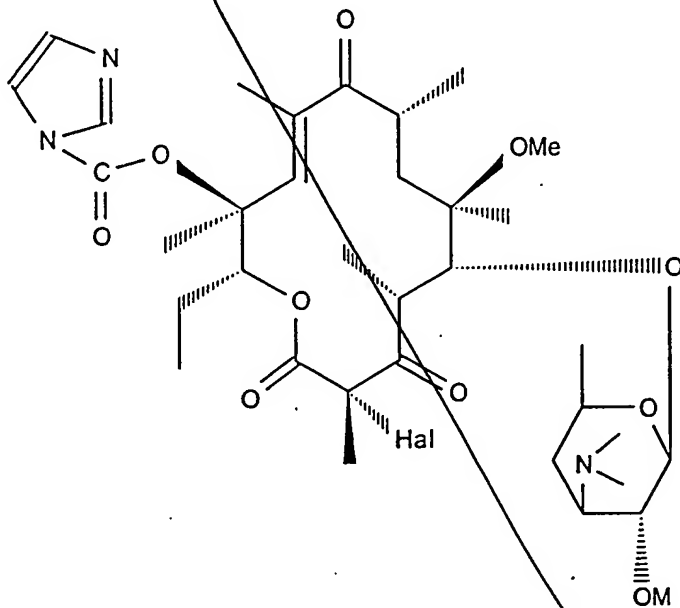
9. An antibiotic composition comprising an antibiotically

effective amount of a compound of claim 7 and an inert pharmaceutical carrier.

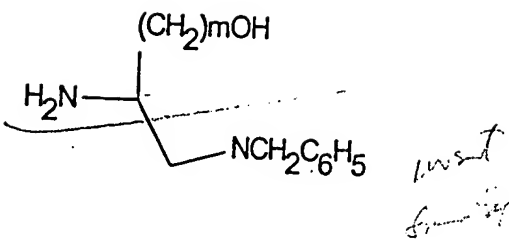
10. A method of treating bacterial infections in warm-blooded
5 animals comprising administering to warm-blooded ^{in need thereof} animals an
antibiotically effective amount of a compound of claim 1.

11. A method of treating bacterial infections in warm-blooded
10 animals comprising administering to warm-blooded ^{in need thereof} animals an
antibiotically effective amount of a compound of claim 7.

12. A process for the preparation of a compound of claim 1
comprising reacting a compound of the formula



wherein Hal is halogen and OM is a protected hydroxyl with a
25 compound of the formula



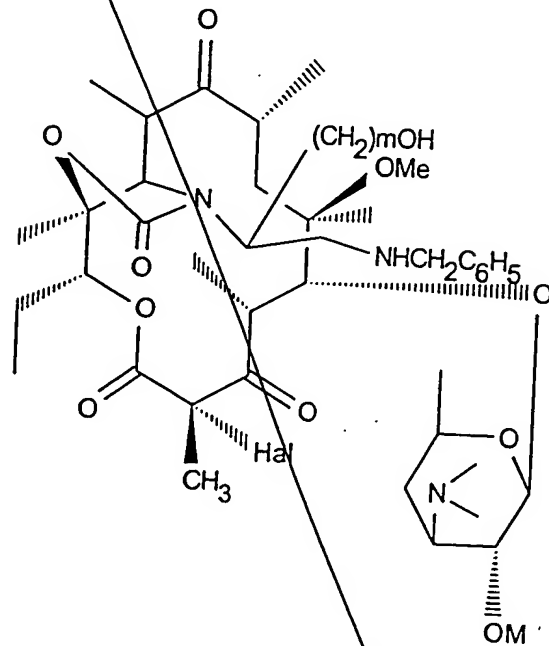
5

Sub

A 6

wherein m is an integer from 1 to 8 to obtain a compound of the formula

10



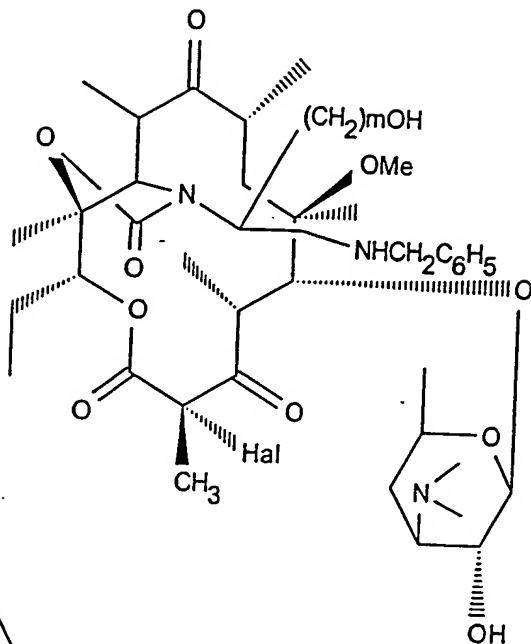
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deprotecting the 2'-hydroxyl to obtain a compound of the formula

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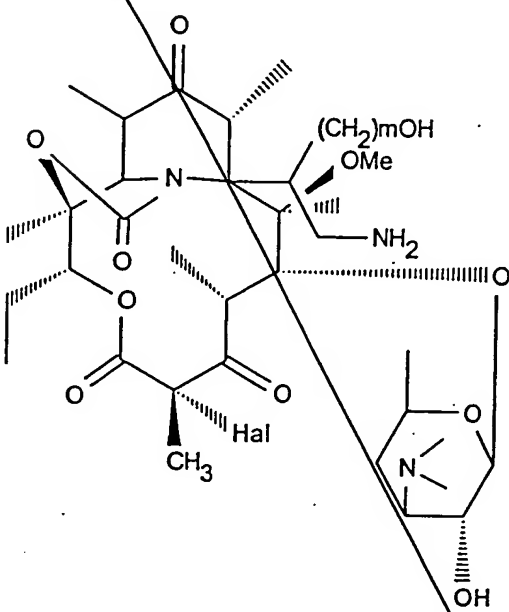
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Sub
A₄

V

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reacting the latter with a debenzylating agent to obtain a compound of the formula

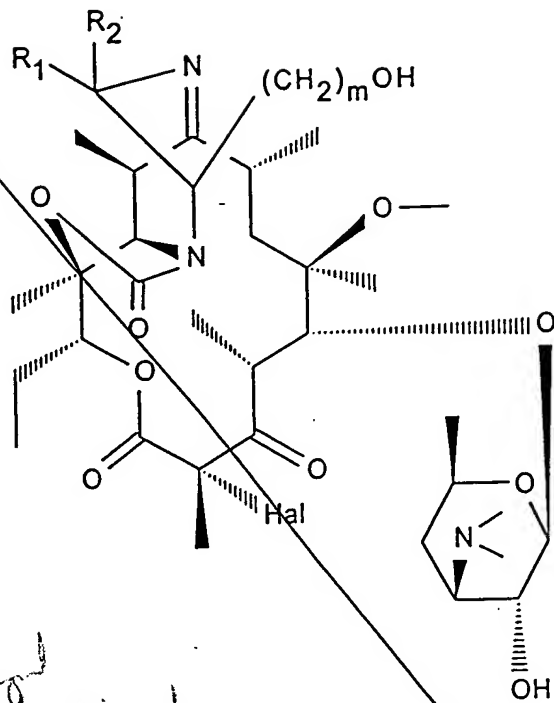


VI

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reacting the latter with a cyclization agent to form a compound of
25 ✓ the formulae

Sub
A6



corresponding to a comp
formula I of claim 1

15 wherein R is -(CH₂)_m-OH and optionally subjecting the latter to an
aralkylating or acylating agent to obtain a compound of claim 1

wherein B is -(CH₂)_n-Ar or -C(=O)-Ar.

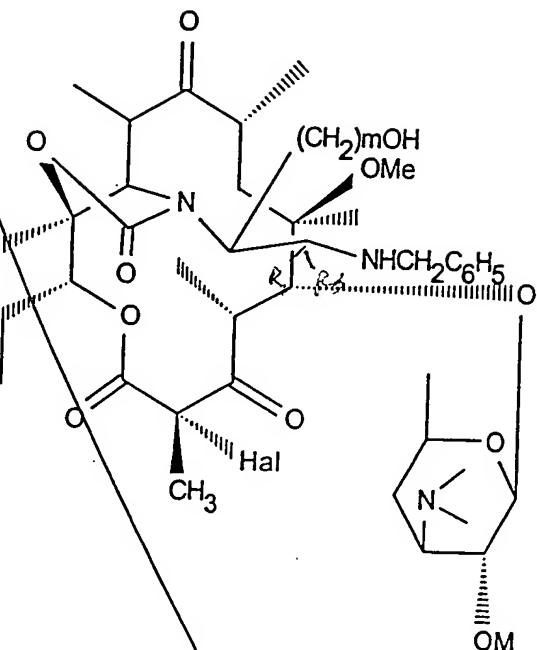
13. A compound selected from the group consisting of

20

Sub
A7

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Sub
A7

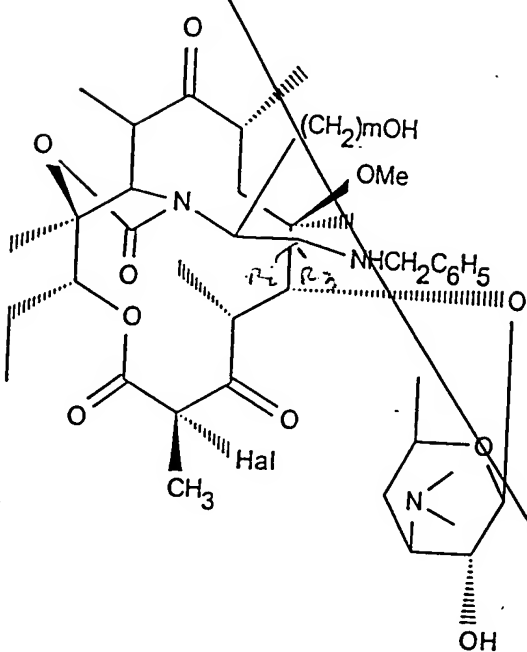


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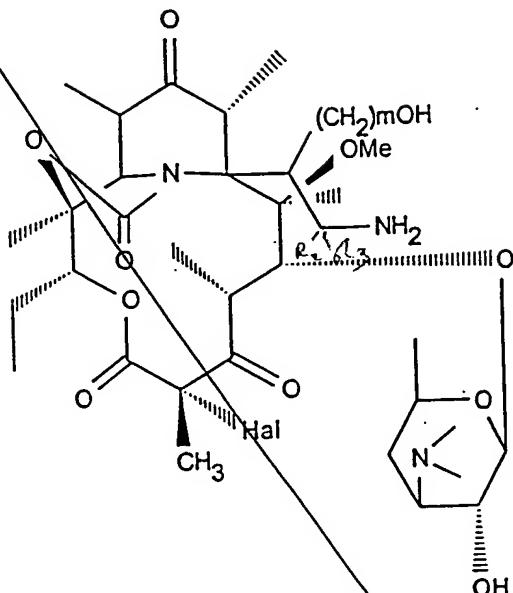
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V



Sub
A7

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VI

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where the substituents are defined as in claim 12.